

EXAMPLE

dissolved in MeOH (300 ml) and a soln, of sodium borohydride (5.02 g) in H<sub>2</sub>O (40 ml) was added dropwise at 0°C over 30 mins., then stirred for 15 mins. Conc. HCl (14.3 ml), satd. NaCl soin. (250 ml) and CH<sub>2</sub>Cl<sub>2</sub> (300 ml) were added to the reaction mixt. The organic layer was fractionated, washed with satd. aq. NaCl soln. (100 ml), dried over anhydrous MgSO<sub>4</sub>, and the solvent was distilled off under reduced press. to give 1-ethoxycarbonyl-3-hydroxypyrrolidine (100 g, 98.7% yield) as an oil.

Followed by prepn. of:
1-ethoxycarbonyl-3-mesyloxypyrrolidine;
1-ethoxycarbonyl-3-phthallmidopyrrolidine;
3-aminopyrrolldine.dihydrochloride; and finally
3-aminopyrrolldine (III).
(4ppW69WSDwgNo0/0).

J61057579-A

86-116676/18 BO3 KANTOH ISHI SEIYAKU KANT- 29,08.84 \*J6 1057-580-A

B(6-D5, 7-D1, 12-A1, 12-D2, 12-G7)

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29.08.84-JP-180212 (24.03.86) A61k-31/39 C07d-205/08 C07d-235 C07d-403/04 C07d-405/04

New 2-azetidinane derivs. - with carcinostatic and antibacterial activity

C86-049841

2-Azetidinone derivs. of formula (1) are new:

R<sub>j</sub> = furyl or methoxyphenyl:

R<sub>1</sub> = benzimidazolyl, <u>phenyl</u>, methoxyphenyl, methoxycarbonylphenyl or ethoxycarbonylphenyl; and

R, = H, phenyl or chloro.

USE

(I) have excellent physiological activity as carcinostatic, immuno-controlling and antibacterial agents and are useful as pharmaceuticals.

PREPARATION

 $R_1 - CH = N - R_2$  (11) • C = C = 0 (111)

STARTING MATERIALS

 $\langle X \rangle$ 

(III) is a reactive and unstable cpd. it is pref. prepd. in situ by treating an acetyl chloride deriv. of formula (V) with an organic amine (IV) (pref. 1-3C alkylamine).

$$R_3 - \begin{pmatrix} 1 \\ C \\ C \\ C \\ C \end{pmatrix} = \begin{pmatrix} C \\ C \\ C \\ C \end{pmatrix} \qquad (III)$$

J61057580-A+

EXAMPLE

A soln. contg. chloroacetylchloride in anhydrous benzene (10 ml) was added dropwise to a soln. contg. (II:  $R_1$  = furyl,  $R_2$  = phenyl) (0.01 mol.) and  $Et_3N$  (1.52 g, 0.015 mol.) in anhydrous benzene (50 ml) at 5-10 °C with stirring. The reaction mixt, was allowed to rise to room temp, and stirred for 2 hrs. The  $Et_3N$ . HCl was removed and the solvent distilled off under reduced press. The residue was chromatographed (silica gel: eluent, hexane-EtOAc) (5: 1-50: 1)) to give (I:  $R_1$  = 2-furyl,  $R_2$  = phenyl,  $R_3$  = II). (8ppW69WSDwgNo0/0).

J61057580- A